=> b reg
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http://www.cas.org/support/stngen/stndoc/properties.html

L1 STR

REP G1=(0-1) AK
VAR G2=C/N
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
GGCAT IS PCY UNS AT 18
DEFAULT ECLEVEL IS LIMITED
ECOUNT IS E1 0 AT 18

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 18

STEREO ATTRIBUTES: NONE

L3 320091 SEA FILE=REGISTRY ABB=ON PLU=ON OC4-C6/ES L5 150 SEA FILE=REGISTRY SUB=L3 SSS FUL L1

100.0% PROCESSED 3308 ITERATIONS 150 ANSWERS SEARCH TIME: 00.00.01

=> b reg
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DICTIONARY FILE UPDATES: 6 JAN 2008 HIGHEST RN 960045-19-6

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REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

=> b hcap FILE 'HCAPLUS' ENTERED AT 16:55:22 ON 07 JAN 2008 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2008 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 7 Jan 2008 VOL 148 ISS 2 FILE LAST UPDATED: 6 Jan 2008 (20080106/ED)

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> d bib abs hitstr 128 tot

128 ANSWER 1 OF 16 RCAPLUS COPYRIGHT 2008 ACS ON STN
AN 2008:1154936 RCAPLUS
THE Method for the production of 5-[4-[4-(5-cyano-3-indolyl]butyl]-1piperalinyl]benrofuran-2-carboxamide
PA Merck Patent G.m.b.H., Germany
OP OFI Int. Appl., 12pp.
CODEN: PIXXO2
P Patent
LA GERMAN
APPROVED
TO PATENT
APPLICATION
TO PATENT NO
TO PATENT

EMN.	CNII																		
	PATENT NO.				KIND DATE			APPLICATION NO.							DATE				
ΡI	W020061	1420	2		A1				2006WO-EP03344						20060412				
	w:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,		
		CN,	CO,	CR.,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,		
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KN,	KP,	KR,		
		KZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,	MW,	MX,		
		MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,		
		SG,	SK,	SL,	SM,	SY,	TJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,		
		VN,	YU,	ZA,	ZM,	ZW													
	RW:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,		
		IS,	IT,	LT,	LU,	LV,	MC,	NL,	PL,	PI,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,		
		CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG,	BW,	GH,		
		GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	BY,		
		KG,	KZ,	MD,	RU,	IJ,	TM												
	DE10200	5019	670		A1		2006	1102		DE 2	005-	1020	0501	9670	2	0050	426		

PRAI DE 2005-102005019670 A DS CASREACT 145:471564

$$\begin{array}{c|c} L & & \\ & & \\ & & \\ & & \\ \end{array} \\ \begin{array}{c} \text{CONH}_2 \\ \text{I} \end{array} \\ \begin{array}{c} \text{CONH}_2 \\ \end{array}$$

S-[4-[4-(5-Cyano-3-indolyi)butyl]-1-piperazinyl]benzofuran-2-carboxamide and/or a physiol. acceptable salt is prepared by the reaction of and/or a physiol. acceptable salt is prepared by the reaction of with 3-(4-piperazin-1-yibutyl)indole-5-carbonitrile in the presence of a Pd-catalyzed coupling using Pd complexes, and/or the formed size converted into an action and salt by treatment with an acid, or by a second method in which a benzofuran-2-carboxamide (II) or an HX addition salt second method in which a benzofuran-2-carboxamide (II) or an HX addition salt acabonitrile, and/or 5-(4-[4-(5-cyano-3-indoly)]-butyl]-1-piperazinyl]benzofuran-2-carboxamide is converted into an acid-addition salt by treatment with an acid.

MINERI (Reactan): SPN (Synthetic preparation): PREP (Preparation): RACI (Reactan): SPN (Synthetic preparation): PREP (Preparation): PREP

ANSWER 2 OF 16 HCAPLUS COPYRIGHT 2008 ACS on SIN 2005:578527 HCAPLUS 143:126601 Effect of vilarodene on 5-HT efflux and re-uptake in the quinea-pig dorsal raphe nucleus Roberts, Claire; Hagan, Jim J.; Bartoszyk, Gerd D.; Kew, James N. C.

CS SO

raphe nucleus
Roberts, Claire, Hagan, Jim J.; Bartosryk, Gerd D.; Kew, James
Roberts, Claire, Hagan, Jim J.; Bartosryk, Gerd D.; Kew, James
Roberts, Claire, Hagan, Jim J.; Bartosryk, Gerd D.; Kew, James
Psychiatry CEDD. GlaxoGmithKline, Harlow, Essex, CM19 5AW, UK
European Journal of Pharmacology (2005), 517(1-2), 59-63
CODEN: EJPHAR; ISSN: 0014-2999
Elsevier B.;
Journal
English

RE.CNT 20 THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

128 ANSWER 1 OF 16 HCAPLUS COPYRIGHT 2008 ACS on STN (Continued)

RE.CNT 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 3 OF 16 HCAPLUS COPYRIGHT 2008 ACS on STN 2005:184699 HCAPLUS 112:123699 HCAPLUS 112:123699 HCAPLUS 112:123699 HCAPLUS 112:123699 HCAPLUS 112:123699 HCAPLUS 112:123699 HCAPLUS 12:123699 HCAPLUS 12:12369 HCAPLUS 12:12369

RE.CNT 41 THERE ARE 41 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

AN DN TI

ANSMER & OF 16 HCAPLUS COPYRIGHT 2008 ACS on SIN
2004:1154699 HCAPLUS
112:32866 no f. indolylbutylpiperazinylbenzofurencarboxamides as serotonin
receptor ligands and/or serotonin reuprake inhibitors
Heinrich, Timo; Boettcher, Henning; Schiemann, Kai;
Hoelremann, Guenter; Van Amssterdam, Christoph; Bartoszyk,
Gerd; Leibrock, Joachim; Seyfried, Christoph
Merck Patent GmbH, German
PCT Int. Appl., 43 pp.
COMEN; PIXXO2
German
CHI 1 IN

KIND DATE APPLICATION NO. DATE

$$\mathbb{R}^{1} \xrightarrow{\mathbb{R}^{2}} \mathbb{N}$$

Title compds. [I, X = N, CH; Rl-R3 = OH, OA, cyano, halo, COR4, CH2R4; R4 = OH, OA, NH2, NHB, NH2; O = CH2, CO, CH; A, B = alkyl, alkovy, alkenyl, alkoxyl, in B = 0.4; dotted line = optional double bondl, were prepared Thus, 5-[4-(4-(5-cyano-)-indolyl) butyl]-1-piperarinyl]benrofuran-2-carboxanide in Me250 was treated dropwise with concentrate RCl under ice cooling followed by stirring for 10 h to give 5-[4-(4-(5-cyano-2-cxo-2,3-dihydro-Haindol-3-yl) butyl]-1-piperarinyl]benrofuran-2-carboxanide as the dihydrochloride. The latter showed 5-HTIA receptor binding activity with ICSO = 2.9 nM. I are useful as anxiolytics, antidepressants, neuroleptics, antihypetrensives and/or for pos. influencing obsessive-compulsive antihypetrensives and/or for pos. influencing obsessive-compulsive age-related memory defects, eating disorders such as bulinia, ans/or sexual dysfunction.

ANSWER 4 OF 16 HCAPLUS COPYRIGHT 2008 ACS on SIN

816438-37-6 RCAPLUS
2-Benzofurancarboxamide, 5-[4-[4-(5-fluoro-2,3-dihydro-2-oxo-1H-indol-3-yl]buty]]-lp-piperarinyl]-, dihydrochloride (9CI) (CA INDEX NAME)

816438-39-8 HCAPLUS
2-Benzofurancarboxamide, 5-[4-[4-(5-cyano-6-hydroxy-1H-indol-3-y1]buty1]-1-piperatinyl]-, hydrochloride (9CI) (CA INDEX NAME)

●x HCl

816438-41-2 HCAPLUS 2-Benrofurancarboxamide, 5-[4-[4-(5-cyano-2,3-dihydro-1H-indol-3-yl]butyl]-1-piperarinyl]-, monhydrochloride (9CI) (CA INDEX NAME)

163521-12-8 714950-88-6 765935-80-6
RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation of indolybutylpiperatinylbenrofurancarboxamides as serotonin
receptor ligands or reuptake inhibitors)
163521-12-8 MCAPLUS
2-Benrofurancarboxamide, 5-|4-|4-(5-cyano-1H-indol-3-yl)butyl)-1piperatinyl) (CA TNDEX NAME)

(Uses)

(prepr. of indolylbutylpiperazinylbenrofurancarboxanides as serotonin receptor ligands or reuptake inhibitors)
714890-70-6 HCAPLUS
2-Benrofurancarboxanide, 5-[4-[4-(5-cyano-6-hydroxy-1H-indol-3-yl]butyl]-1-piperazinyl]- (CA INDEX NAME)

816438-30-9 HCAPLUS 2-BenzofurancarDoxamide, 5-[4-[4-(5-cyano-2,3-dihydro-1H-indol-3-yl]butyl]-1-piperazinyl]- (CA INDEX NAME)

816438-33-2 HCAPLUS 2-Benzofurancarboxamide, 5-[4-[4-(5-cyano-2,3-dihydro-2-oxo-lH-indol-3-yl|buty|]-i-p-pterarinyl|- (CA INDEX NAME)

816438-35-4 HCAPLUS
2-Benzofurancarboxamide, S-[4-[4-(5-cyano-2,3-dihydro-2-oxo-1H-indol-3-y])hutyl]-l-piperazinyl]-, dihydrochloride (9CI) (CA INDEX NAME)

●2 HC1

L28 ANSWER 4 OF 16 HCAPLUS COPYRIGHT 2008 ACS on SIN

714950-88-6 HCAPLUS 2-Bensofurancarboxamide, 5-[4-[4-(5-cyano-6-[(methylsulfonyl)oxy]-1H-indol-3-yl]butyl]-1-ptperazinyl]- (CA INDEX NAME)

765935-80-6 HCAPLUS
2-Benzofurancarboxamide, 5-[4-[4-(5-fluoro-lH-indol-3-yl)butyl]-l-piperazinyl)- (CA INDEX NAME)

RE.CNT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT L28 AN DN TI

ANSMER 5 OF 16 HCAPLUS COPYRIGHT 2008 ACS on SIN 2004:1154698 HCAPLUS 142:93855 Preparation of indolylbutylpiperarinylbenzofurancarboxamides as serotonin receptor indicated inhibitors and/or serotonin receptor ligands. Heinrich, Timo; Boetcher, Henning, Edulemann, Kal; General Capture, Control Capture, Control Capture, Control Capture, Control Capture, Control Capture, Control Capture, C IN

LA	German																	
211111	PATENT :															ATE		
PI	W020041															0040	524	
	W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI,	
										RU,								
										US,								
	RW:									SD,								
										AT,								
										IT,								
					BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,	
	DE103		TD,															
	AU20042									2003						0040		
	CA25									2004						0040		
	EP16															0040		
										GR.								
	Λ.									BG.						pro,	P.I.	
	BR20040									2004						0000	E 2.4	
										CN 2								
	CN18 JP20065	2770	Š		- T		2006	1207		2006								
	MX2005P.	a135	37		à		2006	0309										
	US20061																215 <	:
PRAI	2003DE-														_			
	2004WO-																	
os	MARPAT	142:	9385	S														
GI																		

Title compds. [I; X = N, CH; Rl, R3 = H, OH, OA, cyano, halo, COR4, CH2R4; R2 = H, (halo-substituted) alkyl, alkylaryl, alkylhetercaryl, hetercaryl; R4 = OH, OA, NH2, NHB, NB2; A, B = alkyl; m = 2-6; n = 0-4], were prepared Thus, 3-(4-chlorobutyl)-lH-indole-5-carbonitrile in THF was added to NaH in THF followed by stirring for 30 min, addition of Mer in THF, and stirring with S-(plyperarin-1-yl)-bencofuran-2-carboxamide and E-2N in N-methylpyrrolidine at 120° for 4 hot ogive 5-(4-4-(5-cyano)-1-methyl-lH-indol-3-yl)-butyl]piperarin-1-yl]-bencofuran-2-carboxamide. The latter showed serotomin reuprake innibitory activity with ICS0 = 2.6 nM. I are useful as anxiolytics, antidepresants, neuroleptics, latter showed serotomin reuprake innibitory activity with ICS0 = 2.6 nM. I are useful as anxiolytics, antidepresants, neuroleptics, sleep disorderes, tardive dyskinesia, learning disorders, geriatric memory loss, bulimia, irritable bowel syndrome, and sexual

ANSWER 5 OF 16 HCAPLUS COPYRIGHT 2008 ACS on STN (Continued) 2-Benrofurancarboxamide, 5-[4-[4-(5-cyano-1-propyl-1H-indol-3-y1)butyl|-1-piperacinyl)- (CA INEEX NAME)

816429-19-3 HCAPLUS
2-Benrofuranceaboxamide, 5-[4-(4-[5-cyano-1-(2-pyridinylmethyl)-1H-indol-3-yl|butyl]-1-piperarinyl|- (CA INDEX NAME)

816429-20-6 HCAPLUS
2-Benzofurancarboxamide, 5-[4-[4-(5-cyano-1-(2-phenylethyl)-1H-indol-3-y]|butyl]-1p-piperazinyl]- (CA INDEX NAME)

IT

816429-21-7P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)
(preparation of indolylbutylpiperazinylbenrofurancarboxamides as serotonin
reuptake inhibitors and/or serotonin receptor ligands)
816429-21-7 RCPPLUS
2-Benrofurancarboxamide, 5-[4-[4-6-cyano-1-methyl-1H-indol-3-yl)butyl]-1piperazinyl-, dihydrochloride (9CI) (CA INDEX NAME)

$$\underset{H_2N-C}{\text{N-}} \circ \overset{\text{N-}}{\underset{\text{Ne}}{\bigcap}} \circ \overset{\text{CH}_2)}{\underset{\text{Ne}}{\bigcap}} \circ \overset{\text{C}}{\underset{\text{Ne}}{\bigcap}} \circ \overset{\text{C}}{\underset{\text{Ne}}{\bigcap}$$

RE.CNT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT 128

ANSWER 5 OF 16 HCAPLUS COPYRIGHT 2008 ACS on SIN (Continued)
dysfunction.

1816429-14-8P 816429-15-9P 816429-16-0P
816429-17-1P 816429-18-2P 816429-19-3P
816429-20-P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)
sined compound; preparation of indolylbutylpiperatinylbenzofurancerboxamide
s as serotonin reuptake inhibitors and/or serotonin receptor ligands)
816429-14-8 MCAPLUS
2-Benzofurancarboxamide, 5-14-[4-(5-cyano-1-methyl-1H-indol-3-yl)butyl]-1piperatinyl)- (CA INDEX NAME)

816429-15-9 HCAPLUS
2-Benrofurancarboxamide, 5-[4-[4-(5-cyano-1-ethyl-1H-indol-3-yl)butyl]-1-piperazinyl)- (CA INDEX NAME)

816429-16-0 HCAPLUS
2-Benzofurancarboxamide, 5-[4-[4-(5-cyano-1-(1-methylethyl)-1H-indol-3-yl]butyl]-1-piperazinyl]- (CA INDEX NAME)

816429-17-1 HCAPLUS 2-Benzofurancarboxamide, S-[4-[4-[5-cyano-1-(phenylmethyl)-1H-indol-3-yl]butyl]-l-piperazinyl]- (CA INDEX NAME)

$$\mathsf{H}_2\mathsf{N} - \mathsf{C}\mathsf{N} = \mathsf{CH}_2 \mathsf{D} \mathsf{A} - \mathsf{C}\mathsf{N}$$

RN 816429-18-2 HCAPLUS

L28 ANSWER 5 OF 16 HCAPLUS COPYRIGHT 2008 ACS on SIN

ANSMER 6 OF 16 HCAPLUS COPYRIGHI 2008 ACS on SIN 2004:892421 HCAPLUS COPYRIGHI 2008 ACS on SIN 2004:892421 HCAPLUS Effects of Systemic injections of Vilardone, a selective serotonin reuptake inhibitor and serotonin 1A receptor agonist, on anxiety induced by predator stress in rats Adamec, Robert; Bartossyk, Gerd D.; Burton, Paul Department of Psychology, Memorital University, St. John's, AlB 3X9, Can. European Journal of Pharmacology (2004), 504(1-2), 65-77 CUDEN; ADPHAS: ISSN: 0014-2999 Journal 3

Combine numbers, 150NN 0014-2999
Clasvier B.V.
Journal
English
We examined the effect of Vilarodone, a selective serotonin reuptake
inhibitor (SSRI) and serotonin 1A (5-NIA) receptor agonist [Bartosryk,
inhibitor (SSRI) and serotonin 1A (5-NIA) receptor agonistic properties.
Eur. J. Pharmacol. 322, 147-153), on change in affect following predator
stress. Vilarodone and vehicle injection (i.p.) occurred either 10 min
after predator stress (prophylactic testing), or 90 min prior to
testing). Predator stress involved unprotected exposure of rats to a
domestic cat. Behavioral effects of stress were evaluated with hole
board, plus-mare, and acoustic startle tests 1 wk after stress. Predator
stress increased anxiety-like behavior in the plus-mare and elevated
response to acoustic startle. In prophylactic testing, Vilarodone
Vilarodone increased stress elevation of startle at 10 mg/kg, Higher
doses of Vilarodone (20 and 40 mg/kg) blocked stress potentiation of
startle. In contrast, Vilarodone had no effect on stress potentiation of
anxiety in the plus-mare. In therapeutic testing, Vilarodone increased
Vilarodone had no effect on stress potentiation of anxiety in the plus-mare. The therapeutic testing, Vilarodone increased
Vilarodone had no effect on stress potentiation of anxiety in the
plus-mare. Taken together, the data suggest a prophylactic potential for
Vilarodone in the treatment of changes in hypervigilance following severe
stress.

Vilazodome in the treatment of changes in hypervigilance following seve stress.
183521-12-8, Vilazodome
RI: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
(Biological study); USES (Uses)
(effects of SSRI and servicent in A receptor agonist, Vilazodome, on
183521-22-8 HCABLUS
- RICARUS (SSRI AND STREAM STREAM)
- RICARUS (SSRI AND STREAM STREAM)
- RICARUS (SSRI AND STREAM STREAM STREAM)
- POPPER STREAM STREA

RE.CNT 37 THERE ARE 37 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L28 ANSWER 7 OF 16 HCAPLUS COPYRIGHT 2008 ACS on STN

765935-80-6P
RL: PRC (Pharmacological activity); SPN (Synthetic preparation); BIOL (Siclogical study): PREP (Preparation)
(preparation of [[(indolyl) buty]|piperasinyl]) bencofurancerboxamide derivative and study of its activity as S-HTIA receptor agonist and serotonin re-uptake inhibitor|
765935-80-6 HCAPLUS
2-Bentofurancarboxamide, 5-[4-(4-(5-fluoro-1H-indol-3-yl)butyl]-1-piperarinyl]- (CA INDEX NAME)

RE.CNT 49 THERE ARE 49 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSMER 7 OF 16 HCAPLUS COPYRIGHT 2008 ACS on STN
2004:641081 HCAPLUS
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English CASREACT 141:314299

Systematic structural modifications of [(indoly1)alky1](pheny1)piperarines led to improved selectivity and affinity within this class of 5-HTIA receptor agonists. Introduction of electron-withdrawing groups in position 5 on the indole group raises serotonin transporter affinity, and the cyano group proved to be the best substituent here. S-Fluoro and 5-Cyano substituted indoles show comparable results in in vitro and in the cyano group proved to be the best substituent here. S-Fluoro and 5-Cyano substituted indoles show comparable results in in vitro and in the cyano group proved to be the best substituent here. S-Fluoro and 5-Cyano substituted indoles show comparable results in in vitro and in the cyano group of the comparable sensitive in the comparable sensitive in the comparable sensitive (p-chloroamphetamine assay) and in vivo (ultrasonic vocalization) tests. Optimization of the arylipperarine moiety indicated that the 5-benrofurany1-2-carboxamide was best suited to increase 5-HT transporter and S-HTIA receptor affinity and to suppress D2 receptor binding sensitive to the comparable sensitive in the

or reagent!

(preparation of ||(cyanoindoly|)buty||piperariny||benzofurancarboxamide derivative and study of its activity as S-HTIA receptor agonist and serotonin re-uptake inhibitor|

163521-12-8 RCAPUIS

2-Benzofurancarboxanide, S-[4-[4-(S-cyano-1H-indol-3-yl|)buty|]-1piperariny||- (CA INDEX NAME)

ANSMER 8 OF 16 HCAPLUS COPYRIGHT 2008 ACS on SIN
2004;346288 HCAPLUS
141:88987
A new synthesis of indole 5-carboxylic acids and 6-hydroxy-indole-5-carboxylic acids in the preparation of an o-hydroxylated metabolite of vilazodome
Heinrich, all pharmaceutical Research, Merck KGaA, Darmstadt,
4233. Germaceutical Research, Merck KGaA, Darmstadt,
Pharmaceutical Research, Pharmaceutical Research, Pharmaceutical Resea

so

English CASREACT 141:88987

A major metabolite of the potential antidepressant vilacodome formed in rat, dog, monkey and human liver microsomes is \$-i4-14-(4-cyano-6-hydrouy-1B-indol-3-yl)butyl|-1-piperatinyl|-2-bencofurancarboxamide (I). For the construction of the salicyl-like substituted indol-a synthesis of camoxirole was adapted using Japp-Klingemann-type Fischer-indole synthesis protocols. The reaction of 4-amino-2-hydroxybenoic acid with synthesis protocols, The reaction of 4-amino-2-hydroxybenoic acid with synthesis protocols. The reaction of 4-amino-2-hydroxybenoic acid with synthesis protocols. The reaction of 4-amino-2-hydroxybenoic acid with synthesis protocols. The synthesis protocols are also synthesis protocols. The synthesis protocol. Punctional group interconversion of carboxylic acid via carboxamide into 3-14-(3-6-ind)vac-4-pendyn-1-(2B)-pyridinyl)butyl]-B-indole-3-butanoic acid. Finds-4-by synthesis protocol. 71650-68-69

71650-68-69

(Reactant or reagent) (SPN (Synthetic preparation); PREP (Preparation); PACT (Reactant or reagent) (Castoxy)-hydroxy-Harindole-3-butanotae from (Icarboxy)-chholisyo-Harindole-3-butanotae from (I IT

128 ANSWER 8 OF 16 HCAPLUS COPYRIGHT 2008 ACS on STN (Continued)

163521-12-8DP, Vilazodone, metabolites
RE: SPM (Synthetic preparation); PREP (Preparation)
(preparation of vilazodone metabolite via Japp-Klingemann-type Fischer
indole synthesis of 2.5-dicarboxy-6-hydroxy-1M-indole-3-butanoate from
[[carboxy(schinsy=arboxy)]pentylidenen|hydrazin)(|hydroxy)|benroate

[[Garboxycarbonyv1pentyltdene]]
163521-12-8 HCAPUUS
2-Benzofuracarboxanide, 5-[4-[4-(5-cyano-1H-indol-3-y1]]butyl]-1piperazinyl)- (CA INDEX NAME)

IT

714950-70-6P, 5-[4-[4-(5-Cyano-6-hydroxy-lH-indol-3-yl)butyl]-1piperazinyl]-2-benrofurancarboxamide
RI SPN (Synthetic preparation); PREP (Preparation)
Japp-Klingmann-type Fischer indole synthesis of 2,5-dicarboxy-6hydroxy-lH-indole-3-butanoate from [[carboxy(ethoxycarbonyl)pentylidene]
hydratino|(hydroxy)benroate intermediate)
714950-70-6 RCAPLUS
2-Benrofurancarboxamide, 5-[4-[4-(5-cyano-6-hydroxy-lH-indol-3-yl]butyl]-1piperazinyl)- (CA INDEX NAME)

THERE ARE 25 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L28 ANSWER 9 OF 16 HCAPLUS COPYRIGHT 2008 ACS on STN

AN DN TI

ANSWER 9 OF 16 HCAPLUS COPYRIGHT 2008 ACS on SIN 2002:97808 HCAPLUS 138:46\*71 Polymorphic forms of 1-'4-(5-cyanoindol-3-y))butyl-4-(2-cartamoylbenzofuran-5-y))piperatine hydrochloride Bathe, Andreas; Helfert, Bernd; Neuenfeld, Steffen; Kniel, Heike; Bartels, Matthias; Rudolph, Susanne; Boettcher, Henning Matthias; Rudolph, Susanne; Boettcher, Henning Matthias; Rudolph, Germany COT Inc. 101, Germany CODEN: PIXXX2 Patent English CONT 1 IN

PA SO

	CNT 1				KIND		DATE			NDD:	TCAT	TON:	NO.		DATE			
	PAIGNI							APPLICATION NO.										
PI				20021227 20030220		2002WO-EP06153						20020605						
	w:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,	
		co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,	
		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	IN,	TR,	TT,	TZ,	
		UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZM,	ZW								
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AI,	BE,	CH,	
		CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PI,	SE,	TR,	
		BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG	
	CA24							1227								0020	605	
	AU2002320822									2002.	AU-0.	3208	22		21	0020	605	
	AU20023																	
	EP1397357				A2	20040317												
	R:							FR,				LI,	LU,	NL,	SE,	MC,	PT,	
								MK,										
	EE-2004							0415								0020		
	HU20040															0020		
	CN15															0020		
	BR20020							0817								0020		
	JP20045				I			1118								0020		
	NZ5							0929								0020		
	RU2303598															0020		
	MX2003PA11723															0031		
	US20041							0729										
	IN2004KN00031							0407								0040		

US2004197528 Al 20040729 2003US-0481270 20031319
IN2004NN0031 A 20060407 2004TA-NN0031 20040109
ZA2004000329 A 20050415 2004ZA-0000329 20040115
PRAI 2001EP-0113663 A 20050415 2004ZA-0000329 20040115
AB The invention relates to new crystalline modifications of the hydrochioride salt of 1-[4-(5-cyanonidool-3-yl) butyl]-4-(2-carbanoyl-benofuran-5-yl)-piperazine, crystalline modification of the dihydrochioride of 1-[4-(5-cyanonidool-3-yl) butyl]-4-(2-carbanoyl-benofuran-5-yl)-piperazine, crystalline modification of the dihydrochioride of 1-[4-(5-cyanonidol-3-yl) butyl]-4-(2-carbamoyl-benofuran-5-yl)-piperazine and amorphous 1-[4-(5-cyanonidol-3-yl) butyl]-4-(2-carbamoyl-benofuran-5-yolid pharmaceuticals for the treatment of prevention of depressive disorders, anxiety disorders, bipolar disorders, manla, dementia, substance-related disorders, benezual dysfunctions, eating disorders, obesity, fibromyaljua, sleeping disorders, psychiatric disorders, cerebral infarction, tension, for the threspy of side-effects in the treatment of puerperal lactation. Thus, to a solution of 1-[4-(5-cyanonidol-3-yl)] butyl]-4-(2-carbamoylbenofuran-5-yl)]piperazine in Tiff was added RiCl. The I hydrate obtained was dried at 85-90° to give I which was characterized by spectral properties.

II National Pacatanti; TMU (Therapeutic use); BIOL (Biological study); RACI (Reactant); TMU (Therapeutic use); BIOL (Biological study); RACI (Reactant); CASE (Uses) (preparation of polymorphic forms of (cyanonindolyl))butylcarbamoylbenofurany lipiperazine) (CA: IDEX: NNMS)

128 ARSWER 10 OF 16 HCAPLUS COPYRIGHT 2008 ACS ON STN
AN 2002:391537 HCAPLUS
DN 136:380124 TI Veterinary use of combined 5-HTLA againsts and serotonin reuptake representation for the treatment of traumatic and compulsive disorders associated with behavioral stressors
N Bartossyk, Gerd
PA Merck Patent Gnubh, Germany
D CT Int. Appl., 20 pp.
CODEN: PIXXD2
T Patent
LA English
FANLCNI I
PATENT NO. KIND DATE APPLICATION NO. DATE 

IT

128 ANSWER 10 OF 16 HCAPLUS COPYRIGHT 2008 ACS on STN (Continued)

RE.CNT 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L28 ANSMER 11 OF 16 HCAPLUS COPYRIGHT 2008 ACS on SIN (Continued)
(Biological study); USES (Uses)
(Combined S-HITL agonists and selective serotonin reuptake inhibitors
as analgesics)
RN 163521-12-8 RCAPLUS
CT 2-Beniofurancarboxamide, 5-[4-(4-(5-cyano-1H-indol-3-yl|butyl]-1pipresinyl)- (CA INDEX INDEX

RE.CNT 13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L28 ANSMER 11 OF 16 HCAPLUS COPYRIGHT 2008 AC5 on STN
AN 2002:991504 HCAPLUS
DN 136:380120
TI Novel use of combined 5-HTla agonists and selective serotonin reuptake inhibitors
IN Bartoszyk, Gerd; Sedman, Ewen
DA Merck Patent Gmbh, Germany
SO PCT Int. Appl., 34 pp.
TOT INT. Appl., 34 pp.
TOT PATENT
LA English
FAN.CNT 1
PATENT NO. KIND DATE APPLICATION NO. DATE DT | Datemin | Date | D BR2001015434 JP2004519916 HU2004000504 HU2004000504 CN---1541093 RU---2302243 MX2003PR04341 MX2003PR04341 US2004014771 IN2003RN00778 ZA200304757 2000EP-0125409 2001W0-EP12686

$$\mathsf{NC} = \mathsf{CH}_2 \left[ \mathsf{CH}_2 \right]_{\mathsf{CH}_2}^{\mathsf{H}} = \mathsf{CH}_2$$

The present invention relates to the use of compds. being combined selective serotonin (5-HT) reuptake inhibitors (55RIs) and 5-HTlA receptor agonists, in particular of for a physiol. acceptable salt thereof or 3-[4-[4-(4-cyanophenyl]piperazin-1-y]lbuty]]-HH-indole-5-carbonitrile or a physiol. acceptable salt thereof, for the manufacture of a medicament for the treatment of chronic pain disorders or in treating other conditions where there is hyper-sensitiration to painful signals, hyperalgesia, allodynia, enhanced pain perception, and enhanced memory of pain, as well as for the treatment of irritable bowel syndrome (IBS). T-RCI reduced writhing in mice at 30 mg/kg orally by 82% in pain-relieving acute analgetic property tests. AB tests.
163521-12-8
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL

AU

ANSMER 12 OF 16 HCAPLUS COPYRIGHT 2008 ACS on STN 2001:84453 HCAPLUS 135:228-22 Studies comparing in vivo:in vitro metabolism of three pharmaceutical compounds in rats, dogs, monkeys, and humans [by] using cryopreserved hepatocytes, microsomes, and collages-gel-immobilized hepatocyte cultures Matternative and the particular compounds in the particular compounds in the particular compounds and the particular compounds a

RE.CNT 29 THERE ARE 29 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

AN DN TI

ANGMER 13 OF 16 HCAPLUS COPYRIGHT 2008 ACS on SIN 2001:164199 HCAPLUS 135:441 Systemic EMD 68843 injections reduce anxiety in the shock-probe, but not the plus-maxe test Treit, D., Degroot, A.; Rashluba, S.; Bartoszyk, G. D. Department of Psychology, University of Alberta, Edmonton, AB, T6G 289, Department of Psychology, University of Alberta, Edmonton, AB, T6G 289, European Journal of Pharmacology (2001), 414(2/3), 245-248 CODEN: LSDHAZ; ISSN: 0014-2999 Elsevier Science B.V.

so

CODEN: SJPHAE; ISSN: 0014-2999
Elsevier Science B.V.

Journal

Selective Science B.V.

Journal

Selective serotonin (5-hydroxytryptamine; 5-HT) reuptake inhibitors and
5-HTIA receptor agonists are believed to reduce anxiety. In the present
study we examined the effects of injections of 5-16-14-(5-cyano-1-indoly)butyl-1-piperarinyl-benrofuran-2-carboxamide hydrochloride (BMD 68843),
a 5-HTIA receptor agonist and selective 5-HT reuptake inhibitor, in two
animal models of anxiety, plus-mare and shock-probe. Mats received i.p.

mg/kg) 1 n prior to tearing. Diacepam at the single dose teated and EMD
68843 dose-dependently (significantly at 20 and 40 mg/kg) reduced burying
in shock-probe. However, only diacepam significantly increased open arm
exploration in the plus-maze. Therefore, EMD 68843 has task specific
michiginal control of the second seco

(Uses) (systemic EMD 68843 injections reduce anxiety in shock-probe, but not plus-maze test) (163521-12-8 HCAPLUS 2-Bensofuranearboxamide, 5-[4-[4-(5-cyano-1H-indol-3-yl]butyl]-1-piperainly]- (CA INDEX NAME)

RE.CNT 23 THERE ARE 23 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 14 OF 16 NCAPLUS COPYRIGHT 2008 ACS on STN (Continued)

1-[4-(5-Cyanoindol-3-yl) butyl]-4-(2-carbamoyl-bensofuran-5-yl)-piperarine

1-[4-(5-Cyanoindol-3-yl) butyl]-4-(2-carbamoyl-bensofuran-5-yl)-piperarine

are continued to the stream of the stre

(Uses)
(Compns. of cyanoindolylbutyl(carbanoylbenzofuranyl)-piperazine and its salts for treatment of anxiety and related disorders)
16352-12-8 RCAPLUS
2-Benzofurancerboxanide, S-[4-[4-(S-cyano-1H-indol-2-yl]butyl]-1-piperazinyl)- (CA INDEX NAME)

ANSMER 14 OF 16 HCAPLUS COPYRIGHT 2008 ACS on STN 2000:861478 HCAPLUS 134:32976 HCAPLUS 134:32976 HOvel use of cyanoindolylbutyl(carbamoylbenzofuranyl)-piperatine and its physiologically acceptable salts for treatment of anxiety and related disorders Bartoszyk, Gerd; Seyfried, Christoph; Van Amsterdam, Christoph; Bottcher, Henning; Sedman, Ewen Merck Patent G.m.b.H., Germany PCT Int. Appl., 37 pp. CODEN: PIXXD2

IN

PA SO

	CODEN: Patent English		D2														
	PATENT				KIN		DATE			APPL						ATE	
PI	W020000 W020000	7283 7283	2		A2 A3		2000 2001	1207 1220		20001	WO-E	P043	76		21	0000	516
	W:	ΑE,	AL,														
							GB,										
							KZ,										
							US,						Su,	51,	SK,	SЪ,	IJ,
	DM.	GH.											a.m	20	CH	CV	DP
	1/44						GR,										
			CT.	CM.	GA.	GN.	GW.	MT.	MR.	NE.	SN.	TD.	TG			,	,
	TW5		8		В		2003	0121		TW 1:	999-	8811	9882		1:	9991	115
	CA23	7266	8		A1		2000	1207		2000	CA-2.	3726	68		21	0000	516
	EP11	8527	2		A2		2003 2000 2002 2004	0313		20001	EP-0	9350	31		21	0000	516
	EP11	8527	2		B1		2004	0407									
	R:	AT,	BE,	CH,												MC,	PI,
		IE,	SI,	LT,	LV,	FI,	RO										
	BR20000	11094	R		A		2002	0423		20001	BR-U	0109	48		21	0000	
	CM 13	6160	2		1.2		2002	0321		2001	TK-U	0001	2 C		21	0000	
	RH30030	01103	ć		7.2		2002	0000		20000	CIG-O	0013	22		2	0000	
	HU20020	0127	5		A3		2004	0428		20021	110-0	0012	,,,			5000	310
	JP20035	0044	1		T		2003	0107		2000	JP-0	6209	44		21	0000	516
	AU	77177	8		В2		2004	0401		2000	AU-0	0506	63		21	0000	
	AT2	6356	4		T		2004	0415		20002	AT-0	9350	31		21	0000	516
	BR20000 TR-2003 CN13 HU20020 HU20020 JP20035 AU3 AT2 EP14	1080	0		A1		2004	0421		20041	EP-0	0014	41		21	0000	516
	EP1	1080	0		B1		2006	0823									
	R:								GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
	PT11 RU22 ES23 US65 CZ2 CN16 AT3 EP17	LE,	51,	LI,	LV,	FI,	RO,	0022		2000	n= 0	0250	21		0.	0000	616
	PH25	3747	2		Č2		2004	1010		20001	D11_0	1333	42		21	0000	
	ES22	1934	2		Т3		2004	1201		20001	ES-0	9350	31		21	0000	
	US69	0021	2		В1		2005	0531		2001	US-0	9799	22		21	0000	516
	CZ2	9562	3		В6		2005	0914		2001	CZ-0	0042	26		21	0000	
	CN1€	7957	7		A		2005	1012		CN 2	005-	1005	4417		21	0000	
	AT3	3700	8		T.		2006	0915		2004	AT-0	0014	41		21	0000	
	EP1	3615	6		A2		2000	1227		20061	EP-U	01/2	5 L		21	0000	210
	R:	AT,	BE.	CH.	CY.	DE.	DK.	ES.	FT.	FR.	GB.	GR.	TE.	TT.	LT.	Tall.	MC.
							RO,		,	,	,	,	,	,	,	,	,
	ES22				Т3		2007	0416		20041	ES-4	0014	41		21	0000	516
	NO20010	0574	6		A		2001	1126		2001	NO-0	0057	46		21	0011	126
	NO3	32212	0		В1		2006	0814									
	MX2001E	A121	72		A		2002	0722		2001	MX-P.	A121	72		21	0011	127
	ZA20010	11048	5		A		2003	0630		2001	ZA-U	0104	85		21	0011	220
	HK1(	04844	3.7		Δ1		2005	1209		2001	RK-U	1006	17		21	0030	153
	US20051	1338	6		A1		2005	0526		20041	US-0	9942	26		21	0041	123
	NO20010 NO20010 MX20010 IN20010 HK10 US20051 NO20060 NO3 1999EP- 2000CN- 2000EP- 2004EP-	0156	2		A		2001	1126		20061	NO-0	0015	62		21	0060	406
	NO3	2423	0		B1		2007	0910									
PRAI	1999EP-	0109	295		A		1999	0527									
	2000CN-	-0808	135		A.3		2000	0516									
	2000EP-	0935	031		A3		2000	0516									
	2004EP-	0001	441		A3		2000	0516									

ANSWER 15 OF 16 HCAPLUS COPYRIGHT 2008 ACS on STN 1996:689356 RCAPLUS 125:328501 Preparation of S-aminobenrofuran-2-carboxylates as drug intermediates Bathe, Andreas; Reffert, Bernd; Boettcher, Henning; Schuster, Kurt Merck Patent falbh, German Bernd, Pat. Appl., 13 pp. CODEN: EDEXIM

	CNT 1 PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
	PATENT NO.			APPLICATION NO.	
PI	EP738722	A1	19961023	1996EP-0105701	19960411
	EP738722	B1	20030625		
	R: AI, BE, CH,	DE. DE	. ES. FR.	GB, GR. IE, IT, LI, LU,	NL. PT. SE
	DE19514567	Al	19961024	1995DE-1014567	19950420
	EP1215210			2002EP-0006144	19960411
	EP1215210		20020626		
	EP1215210	B1	20061018		
		DE, DK	, ES, FR,	GB, GR, IT, LI, LU, NL,	SE, PT, IE,
	SI, LT, LV				
	AT243689	T	20030715		19960411
	PI738722	I	20031128		19960411
	ES2201143	T3	20040316	1996ES-0105701	19960411
	AT342893	T	20061115	2002AT-0006144	19960411
	ES2275765		20070616		
	CN1140171	A	19970115		
	AU9650734	A	19961031		19960417
	AU704495	B2	19990422		
	RU2159238	C2	20001120		
	SK284862	B6	20060105		19960417
	SK285224	B6	20060907		
	CA2174494	A1	19961021		19960418
	NO9601579 ZA9603155	A	19961021		19960419
	JP08291161	A A	19961025		19960419
	JP3874837		20070131		19960419
	HU9601033	B2 A2	19971028		19960419
	US5723614	A2	19980303		19960419
	CZ294697	B6	20050216	1996CZ-0001131	
	USS977112	A	19991102	1997US-0960459	
	JP2006290905	A	20061026		
DRAT	1995DE-1014567	A	19950420		20000007
FIGHT	1996EP-0105701	A3	19960420		
	1996JP-0120781	A3	19960419		
	199605-0634825	A 3	19960419		
os	MARPAT 125:328501		1000010		
0.3	PMIENT 123.326301				

Title compds. [I; R = cyano, CO2H, alkoxycarbonyl, etc.; Rl = NH2, piperarino, (N-benyl)piperarinyl, etc.] were prepared Thus, Et 3-carboxylate (preparation described) was converted in 5 1-carbox to 5-piperarinobenofuran-2-carboxanide. 165521-129 [PR | Co2H | C

L28 ANSWER 15 OF 16 HCAPLUS COPYRIGHT 2008 ACS on STN (Continued)

L28 ANSWER 16 OF 16 HCAPLUS COPYRIGHT 2008 ACS on STN (Continued)

1.28 ANGMER 16 OF 16 HCAPLUS COPYRIGHT 2008 ACS on STN
AN 1995:586488 RCAPLUS
N 123:9463
TI Preparation of (indolylalkyl)piperidines and -piperatines as drugs.
Bosetcher, Henning; Seyfried, Christoph; Bartoszyk, Gerd;
Greiner, Hartmut
N Merck Patent G.m.D.H., Germany
COCET, GAXEAN
LOCATE GAXEAN
LA German
LA German

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	DE4333254		19950406	1993DE-4333254	19930930
	EP648767	A1	19950419	1994EP-0114798	19940920
	EP648767	B1	19970528		
				GB, GR, IE, IT, LI, LU,	
	AT153663	T	19970615	1994AT-0114798 1994ES-0114798	19940920
	ES2105454	T3	19971016	1994ES-0114798	19940920
	AU9474244	A	19950413	1994AU-0074244	19940927
	AU679774				
	CN1106811		19950816	1994CN-0116585	19940927
	CN1056610		20000920		
	CA2133152		19950331	1994CA-2133152	19940928
		A1	19950331		
				1994JP-0233538	
				1994PL-0305216	19940928
	CZ293558		20040616		
	NO9403616			1994NO-0003616	19940929
	NO306948	B1	20000117		
				1994ZA-0007622	
	HU71833		19960228	1994HU-0002806	19940929
	HU218918		20001228		
	US5532241				
	RU2132848	C1	19990710		
	5K281793		20010806		
	JP2007119502	A		2007JP-0034671	20070215
PRAI	1993DE-4333254	A			
	1994JP-0233538	A3	19940928		
os GI	MARPAT 123:9463				

AB Title compds. II, X = (No. alkoxy-, cyano-, halo-, R2CO-, R2CO-, R2CO-2-substituted) 3-indoly); R1 = (cyano-, NoCN-2-, alkoxymethy)-, R2CO-2-substituted) 3-indoly); R1 = (cyano-, NoCN-2-, alkoxymethy)-, R2CO-2-substituted) beneforana-6-2 = (cyano-, NoCN-2-, alkoxymethy)-, R2CO-2-substituted) beneforana-6-2; R2CO-2-, R2CO-2

=> d bib abs hitstr 129 tot

NC NH2

129 ANSWER 1 OF 17 HCAPLUS COPYRIGHT 2008 ACS on STN (Continued)

RE.CNT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

RE.CNT 7

ANGMER 4 OF 17 HCAPLUS COPYRIGHT 2008 ACS ON STN
2005:547557 HCAPLUS
143:53543
The combination of a serotomin reuptake inhibitor and a histamine 3 receptor antagonist, inverse agonist or partial agonist, and therapeutic use thereof
use thereof
H. Bundbeck A/S. Den.
PCT Int. Appl., 36 pp.
CODEN: PIXEND
Datent
English
CHAPLES APPLICATION NO. DATE AN DN TI IN PA SO PRAI

ANSWER S OF 17 HCAPLUS COPYRIGHT 2008 ACS on SIN
AN 2005;471926 HCAPLUS
DN 143186623
COPYRIGHT 2008 ACS on SIN
AN 2005;471926 HCAPLUS
COPYRIGHT 2009
COPYRIG 2004W0-EP13070 W 20041117 CASREACT 143:26625; MARPAT 143:26625

Use of title compds. e.g. [f; Rl = H, alkyl, fluoroalkyl, alkenyl, alkynyl, cycloalkylalkyl, bridged cycloalkyl, etc.; R2 = fluoroalkyl; R3 = alkyl, amino, carboxamide| for preparation of a medicament for treatment of depressive disorders is claimed. Thus, a mixture of 400 Med Treatment of depressive disorders is claimed. Thus, a mixture of 400 Med Treatment of 20 min. with bloome in MeOH followed by heating at 400 for 23 h. AcOH and 5-Me 2-thlopseudourea were added followed by concentration and heating at 110° overnight. AcOH was added and the mixture was cooled to 50° followed by pheating at 50° for 21 h. The mixture was cooled to 50° followed by pheating at 50° for 21 h. The mixture was cooled cool of 20° and aqueous Ma sulfite was added over 20° min. followed by aging for 1 h to give 90% 2-methylsulfonyl-4-44—(methylsulfonyl) phenyl 1-6-trifluoromethylsyrimidine. The latter was (methylsulfonyl) phenyl 1-6-trifluoromethylsyrimidine. The latter was 10° mixture was 4-(4-(methylsulfonyl)) phenyl 1-6-trifluoromethylypyrimidine (II). In the chronic insecapable shock in rats model, II at 10 mg/kg orally with paroxetine 5 mg/kg orally gave a full reversal of the chronic escape deficit.

16352-12-8 mtm 68843
Ri. THU (Therapeutic use); BIOL (Biological study); USES (Uses) (coedministration; preparation of pyridines, pyrimidines, and

129 ANSWER 4 OF 17 HCAPLUS COPYRIGHT 2008 ACS on STN (Continued)

L29 ANSMER S OF 17 HCAPLUS COPYRIGHT 2008 ACS on SIN (Continued)
pyracolopyridazines as cyclooxygenase-2 inhibitors for the treatment of
depressive disorders)
RN 163521-12-8 HCAPLUS
CN 2-Benrofurancarboxamide, 5-[4-[4-(5-cyano-1H-indol-3-yl|butyl]-1piperarinyl)- (CA INDEX NAME)

ANSWER 6 OF 17 HCAPLUS COPYRIGHT 2008 ACS ON STR 2005.179317 HCAPLUS 143:27004 HCAPL AN DN TI IN PA SO | Patent | Ranglish | Column | PRAI IT

L29 ANSWER ? OF 17 HCAPLUS COPYRIGHT 2008 ACS on STR
AN 2005:136555 HCAPLUS
II Selective serotonin reuptake inhibitors for the treatment of prenature female orgasm
IN May, Kathryn Elizabeth; Quinn, Paul
A Pfizer Limited, UK; Dfizer Inc.
50 PCT Int. Appl., 20 pp.
COMBEN PIXAD2
LA English
FAN.CNT 1
PATENT NO. KIND DATE APPLICATION NO. DATE PRAI AB IT

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129 ANSWER 6 OF 17 HCAPLUS COPYRIGHT 2008 ACS on STN (Continued)

RE.CNT 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

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ANSWER 8 OF 17 HCAPLUS COPYRIGHT 2008 ACS on SIN
AN 2004:452952 HCAPLUS
DN 141:1296
IN Method of using a cyclooxygenase 2 (COX-2) inhibitor and a 5-HTIA receptor modulator as a combination therapy for pain, inflammation, and other

N Stephenson, Diane T.; Taylor, Duncan P.
PA Pharmacia Corporation, USA
OPCT Int. Appl., 195 pp.
CODEN: PIXXD2
DT Patent
LA English
PARLENT IN N. KIND DATE APPLICATION NO. DATE
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PB DT LA AB

ANSMER 9 OF 17 HCAPLUS COPYRIGHT 2008 ACS on STN 2004:371879 HCAPLUS 101:368059

Phase I and II entyme characterization of two sources of HepG2 cell lines Phase I and II entyme characterization of two sources of HepG2 cell lines In Vitro Technologies, Baltimore, ND, 21227, USA Xenobiotica (2004), 34(3), 243-256

Xenobiotica (2004), 34(3), 243-256

Iaylor & Francis Ltd.

Journal Logist State St

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L29 ANSWER 10 OF 17 HCAPLUS COPYRIGHT 2008 ACS on SIN

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ANSWER 10 OF 17 HCAPLUS COPTRIGHT 2008 ACS on STN
AN 2004:2708 HCAPLUS
DN 140:33457 septiate inhibitor combination with a GABAB receptor antagonist
for the treatment of depression and other disorders
IN Mork, Arne: Cremers, Thomas Ivo Franciscus Hubert; Willigers, Sandra
A. H. Lundbeck A/S, Den
COODEN: PIXXD2
COODEN: PIXXD2
LA English
FAN.CNT 1
LA English
FAN.CNT 1
PATENT NO. KIND DATE APPLICATION NO. DATE 

2-Benzofurancarboxamide, 5-[4-[4-(5-cyano-1H-indol-3-yl]butyl]-1-piperazinyl]- (CA INDEX NAME)

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L29 AMSWER 11 OF 17 HCADLUS COPYRIGHT 2008 AC5 on STN
NN 2002:1006815 HCADLUS
NN 1002:1006815 HCADLUS
NI 140:135974
TI Treatment for depression and anxiety by the combination of a PDE IV inhibitor and an anxietyresant or an anxiolytic agent
NN 50bolov-Jaynes, Susan Beth; Schmidt, Christopher Joseph
PA Pfizer Products Inc., USA
SO PCT Int. Appl., 62 pp.
TO PARTICLE TAXAD2
LA English
FAN.(NT 1
PATENT NO. KIND DATE APPLICATION NO. DATE
                                                 PI W02003105902
                                                    MX2004PAIR855 A 20050331 200MMC-PAIR855 20041126 <--
TRX004CN03177 A 2006030 2004IN-CN03177 20041213 <--
2002UB-38918IP P 20020617 <--
2002UB-38918IP P 20020617 <--
2003MO-1802295 W 20030605 <--
MARART 140:35974 W 20030605 <--
MARART 140:35974 The present increased. Including a human. Dy administering to the mammal a PDE IV inhibitor in combination with an antidepressant or an anxiolytic agent. It also relates to pharmaceutical compus. containing a pharmaceutically acceptable carrier, a PDE IV inhibitor and an anxiolytic agent of the present of the present of the present of the pharmaceutical compus. Containing a pharmaceutically acceptable carrier, a PDE IV inhibitor and an anxiolytic agent or an anxiolytic agent or antidepressant or an anxiolytic agent of the present of the
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L29 ANSWER 12 OF 17 HCAPLUS COPYRIGHT 2008 ACS on STN

RE.CNI 54 THERE ARE 54 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 12 OF 17 HCAPLUS COPYRIGHT 2008 ACS on STN 2002:659259 RCAPLUS 138:26839 (RAPLUS 138:26839 (RAP

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302(3), 1220-1228
American Society for Pharmacology and Experimental Therapeutics
American Society for Pharmacology and Experimental Therapeutics
American Society for Pharmacology and Experimental Therapeutics
Maglish
EMD 68843 (vilarodone) is a novel compound with combined high affinity and
selectivity for the 5-hydroxytryptamine (5-HT) transporter and 5-HT1A
receptors. EMD 68843 was tested as a prototype compound, which benefits
from dual pharmacol effects that could increase extracellular 5-HT1A
receptors. EMD 68843 was tested as a prototype compound, which benefits
from dual pharmacol effects that could increase extracellular 5-HT1A
receptors as for the second of the full
5-HT1A receptor agonist R-(1)-trans-8-hydroxy-2-(N-n-propyl-N-(19-iodo-29propenyl) jaminoterial in 9-00-FFPAT1, indicating that it is a partial
agonist at 5-HT1A receptors. Acute, systemic administration of EMD 68843
agonist at 5-HT1A receptors. Acute, systemic administration of EMD 68843
receptors agonist at 5-HT1A receptors. The second propenyl layer (19-iodo-29propenyl) jaminoterial in 9-00-FMPAT1, indicating that it is a partial
agonist at 5-HT1A receptors. Acute, systemic administration of EMD 68843
receptors agonist at 5-HT1A receptors. The second propenyl 1-MT1A
agonist at 5-HT1A receptors. The second propenyl 1-MT1A
agonist at 5-HT1A receptor and the second propenyl 1-MT1A
agonist at 5-HT1A receptor of the second propenyl 1-MT1A
agonist at 5-HT1A receptor of the 1-MT1A receptor of the 1-MT1A
agonist at 5-HT1A receptor of 5-HT release in the HDV and KT by 5-HT1A
agonist at 5-HT1A receptor. The second propenyl 1-MT1A
agonist at 5-HT1A receptor of 5-HT release in the HDV and KT by 5-HT1A
agonist at 5-HT1A receptor of 5-HT release in extracellular 5-HT were greatly reduced in the Fortal and mice but only within a narrow dosage range. Like HD 68843 blocked expression of the
5-HT1 behavioral syndrome induced by 8-00-DDAT. Taken together, the
results show that EMD 68843 augments extracellular 5-HT levels in
forebrain regions to a grea

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ENTERIN NO. RIND DATE APPLICATION NO. DATE

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M02001070230
A2 20010927 200100-18055740
WE AN JR. CA. CN. JP, KR. MX, PL, US. 2A.
FM: AT, BE, CH, CY, DE, DK, ES, FT, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, CH. A2 20010212
ED--1267878
R. CA--2399985
A1 20010223 --RI AR BE, CH, GR DE, ES, FR, GB, GR, IT, IL, LU, ML, SE, MC, PT, RE, TR, CY, TR
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R. F. TR
RECOLOGIST AND A 20030603 2001ER-0914448
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D 20000317 --ENGLOSST-04 A 2003031 --ENGLOSST-04 A 2003031 --ENGLOSST-04 A 2003031 --ENGLOSST-04 A 2003031 --ENGLOSST-04 B 20010213 --ENGLOSST-04 B 2001021 --ENGLOSST-04 B 20010213 --ENGLOSST-04 B 2001021 --ENGLOSST-04 B 2001021 --ENGLOSST-04 B 2001021 --ENGLOSST-04 B 20010 PRAI

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129 ANSWER 14 OF 17 HCAPLUS COPYRIGHT 2008 ACS on STN (Continued)

L29 AN DN TI

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Compounds with 5-HTIa agonist activity useful for treating disorders of the outer retina Collier, Robert J., Jr.; Kapin, Michael A.; Hellberg, Mark R.; Dean, Thomas K.

PCT Int. Appl., 23 pp.

CCDERN: PIXXO2

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	RW:	AI,	BE,	CH,	CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL.	
		PT,	SE,	TR														
	CA24				A1		2001			2001						0010		
	EP12	6350	4		A2		2002	1211		2001	EP-0	9182	0.8		2	0010	223	<
	EP12	6350	4		B1		2003	0820										
	R:		BE,			DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PI	
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	ES22	0484	8		Т3		2004	0501		2001	ES-1	9182	0.8		2	0010	223	<
	TW2	6877	7		В		2006	1221		TW 2	001-	9010	6235		2	0010	316	<
	ZA20020	0635	0		A		2003	0808		2002	ZA-0	0063	50		2	0020	808	<
	US20032	0789	0		A1		2003	1106		2002	US-0	2210	70		2	0020	909	<
	KR7	4919	1		B1		2007	0813		2002	KR-0	7121	70		2	0020	916	<
	MX2002P	A090	72		A		2003	0312		2002	MX-P.	A090	72		2	0020	917	<
	HK10	5150	4		A1		2004	0423		2003	HK-0	1034	44		2	0030	515	<
	AU20052	0260	0		A1		2005	0707		2005.	AU-0	2026	0.0		2	0050	615	<
	US20052	5612	9		A1		2005	1117		2005	US-0	1874	74		2	0050	722	<
PRAT	2000015-	1902	79P		P		2000	0317	<-	-								

200005-1902799 P 20000317 <-2001W0-US05700 W 20010223 <-2001W0-US05700 W 20010223 <-2002US-0221070 Al 20020909 <-Compns. and methods are disclosed for treating disorders of the outer
retina with compds. with 5-HTlA agonist activity, e.g. buspirone.
H. BAC (Blood)cal activity or effector, except adverse) RSU (Biological
study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES
(Uses)
(5-HTlA agonist for treating disorder of outer retina)
16321-12-8 RCAPLUS
2-BenioChuBuncarboxamen, 5-[4-[4-(5-cyano-HH-indol-3-yl]butyl]-1piperasinjl-) (CA INDEX NAME)

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AMSWER 15 OF 17 HCABLUS COPYRIGHT 2008 ACS on STN 2001:578501 HCAPLUS 135:1578501 HCAPLUS 135:157861 HCAPLUS 135:157861 HCAPLUS 135:157861 HCAPLUS 135:157861 HCAPLUS 135:157861 Distinct temporal pattern of the effects of the combined serotconin-reuptake inhibitor and S-HT1A agonist EMD 68843 on the sleep EEG in healthy Release R. M.; Attonity-rich; T. A.; Steiger, A. Murck, H.; Friebese, R. M.; Attonity-rich; T. A.; Steiger, A. Murck, H.; Friebese, R. M.; Attonity-rich; T. A.; Steiger, A. Murck, H.; Friebese, R. M.; Attonity-rich; T. A.; Steiger, A. Murck, H.; Friebese, R. M.; Attonity-rich; T. A.; Steiger, A. Murck, H.; Friebese, R. M.; Attonity-rich; T. A.; Attonity-rich;

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AN 2000:605068 HCAPLUS
N 134:292112
TI Drug action at the 5-HTIA receptor in vivo: autoreceptor and postsynaptic receptor occupancy examined with DET and (carbonyl-120 MAY-100835)
AN Sedman, B.; Cowen, P. J.; Grasby, P. M. R.; Bargent, P. A.; Mccaer, E.; Sedman, B.; Cowen, P. J.; Grasby, P. M. R.; Bargent, P. A.; Mccaer, E.; Sedman, B.; Cowen, P. J.; Grasby, P. M. Sedman, P. Sedma

RE.CNT 41 THERE ARE 41 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

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129 ANSWER 17 OF 17 HCAPLUS COPYRIGHT 2008 ACS on STN

AN 2000:98327 KCAPLUS

132:186650

II Treating depression with a combination of a serotonin uptake inhibitor, a 5-HIJA presynaptic antagonist, and a 5-HIJA agonist

IN Depoortere, Henri

N Depoortere, Henr
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RE.CNT 11 THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

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L6 1 US20070099933/PN 1 US20060160824/PN L7

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30 L12 AND (PD<=20040524 OR AD<=20040524 OR PRD<=20040524) L13

L14 2 L12 AND L6-7 E HEINRICH T/AU 23 E3-4

L15 E HEINRICH TIMO/AU

L16 41 E3

E BOTTCHER H/AU

96 E3-6 L17

E BOTTCHER HENNING/AU

L18 9 E3

E SCHIEMANN K/AU 43 E3-4

E HOLZEMANN G/AU

L20 17 E3-5 E VAN AMSTERDAM C/AU

L21

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E BARTOSZYK G/AU 122 E4-8

E LEIBROCK J/AU

44 E3-6 L23

E SEYFRIED C/AU

114 E3-6 E SEYFRIED CHRISTOPH/AU

116 E3-5 T<sub>2</sub>25

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L27 16 L12 AND L15-26

L28 16 L14,L27

17 L13 NOT L28 T<sub>1</sub>29

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